Data analysis standards in metabolomics

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Aims and goals

It is clear that algorithms do not drive metabolomics investigation, but rather the question one seeks to answer with metabolomics dictates the data analysis strategy. The goal of this group is to define the reporting requirements associated with statistical and chemometric analysis of metabolite data. This will include identifying the type of algorithm that will be required, and where a model is built, its construction and its validation. These points must be reported so that the data analysis is as objective and unbiased as possible.

Scene setting

The figure opposite identifies the clear flow of information (pipeline) in a typical metabolomics experiment. Whilst multivariate analysis (MVA; also referred to as chemometrics and machine learning) features at the end of the flow, in order for the analysis to be valid there must be robust experimental design. For MVA this particularly refers to the sample type the numbers needed and obviously using the correct control and test groups. Althour

Robust experi	mental design
Robust and rep	roducible data
Well curated databases	
l Validated data analysis	

obviously using the correct control and test groups. Although experimental data capture and data storage and retrieval are also important, these are dealt with by other working groups.

Design of experiments (DOEs) require that the biological space is adequately populated prior to data capture and subsequent analysis. This is clearly determined by the experiment in question but, for example, if one was interested in the childhood disease leukaemia the control set of healthy individuals must not include adults. Most MVA algorithms are only capable of interpolation, that is to say they give answers within their knowledge realm and can not extrapolate beyond this. Therefore to account for this the DOE would span the metadata that were collected in terms of e.g. sex, age, height, BMI (body mass index) etc, and include suitable sample numbers to account for inherent biological variability. There are approaches to accomplish the former based on space filling algorithms including full or fractional factorial design, Plackett-Burman, Taguchi arrays, to name the most popular ones. The latter requires some preliminary metabolite data collection of the same samples, nominally under identical conditions, where the variation in metabolite data can be assessed in terms of biological reproducibility. Power laws, ANOVA (analysis of variance) and MANOVA (multivariate ANOVA) can then be used to decide on the minimum number of samples required.

Reporting structure:

The number of samples per class should be reported along with the relevant metadata capture, and how accurately these are spanned in the calibration, validation and test sets (*vide infra* for definitions of these data sets).

Pre-processing

Before any analysis is performed metabolite data must be scaled / normalised. There are many approaches that can be used and the most poplar include scaling to total response, scaling to individual metabolite (or peak), log transformation, scaling to unit variance (autoscale), Pareto scaling, derivatisation, mean centring, vector normalisation. The way in which the data were scaled prior to analysis must be explained. In most instances this will have been optimised, and if this is the case then this must be performed objectively as described under validation below.

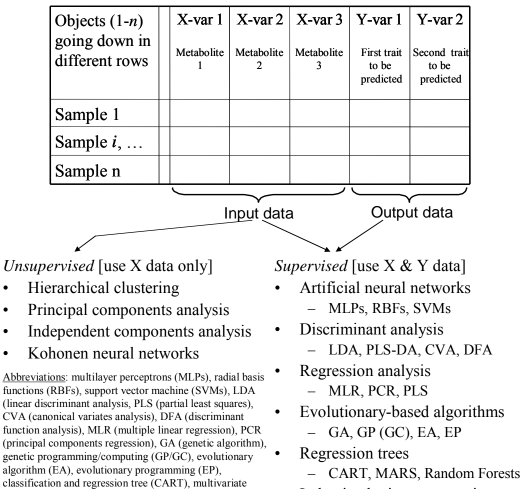
Reporting structure:

The way in which the data are scaled prior to analysis must be explicitly detailed.

Algorithm selection

The sort of question that one wants to answer drives the selection of the most relevant algorithm (or set of algorithms). It is not feasible to discuss the pros and cons of each method as this is often subjective, but we can define a reporting structure based on the biological application.

Multivariate data consist of the results of observations of many different metabolites (variables) for a number of individuals (objects). Each variable may be regarded as constituting a different dimension, such that if there are *n* variables (metabolites) each object may be said to reside at a unique position in an abstract entity referred to as *n*-dimensional hyperspace. This hyperspace is necessarily difficult to visualise, and the underlying theme of multivariate analysis (MVA) is thus *simplification* or dimensionality reduction. This dimensionality reduction occurs in one of two ways; either using an unsupervised or supervised learning algorithms (see the figure below for a summary of the main methods).



Inductive logic programming

Unsupervised learning

adaptive regression splines (MARS).

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When learning is unsupervised, the algorithm is shown a set of inputs and then left to *cluster* the metabolite data into groups. For MVA this optimization procedure is usually *simplification* or dimensionality reduction. This means that a large body of metabolite data (x-data) are summarised by means of a few parameters with minimal loss of information. Typically principal components (PCA) and hierarchical cluster analyses (HCA) are used, and after clustering the ordination plots or dendrograms then have to be interpreted.

Reporting structure:

As PCA and HCA are unbiased analysis and describe the natural variation in the input x-data what needs to be reported is the percent explained variance generated for each principal component plotted and the specific way in which HCA has been generated. This includes construction of the similarity matrix and whether an agglomerative or divisive clustering algorithm is used.

Supervised learning

When one knows the desired responses (y-data, or traits or classes) associated with each of the metabolite data inputs (x-data) then the system may be supervised. The goal is to find a mathematical transformation (model) that will correctly associate all or some of the inputs with the target traits. This trait can be categorical (e.g., disease vs. healthy) or quantitative (e.g., grade of cancer, response to therapy). In its conventional form this is achieved by minimising the error between the known target and the model's response (output). In addition

there exist special types of supervised learning that effect explanatory analyses; that is to say the mathematical transformation from input to output data is transparent. Such inductive methods allow one to discover which metabolites (inputs) are key for the separation of the traits to be predicted. These approaches may help in the validation of the model in terms of its biological relevance that can be tested by a complementary approach using transcriptomics and proteomics.

Validation: As these supervised learning methods use both input x-data and output y-data in model formation the analysis must be fully validated. All these methods require optimisation. For regression based approaches (MLR, PCR, PLS) and discriminant analysis (LDA, CVA and DFA) the number of latent variables that are used in the model must be optimised. For neural networks the optimisation will be in terms of the number of iterations (epochs) in model formation, and for evolutionary-based algorithms the number of population cycles. Whilst for pre-processing, the effect on model performance must also be assessed objectively.

Leave-one or leave-n out for model calibration is where a single or n sample(s) are iteratively left out, the model is then reconstructed, the omitted sample projected into model space, and its location used to assess the predictive ability of the model. However, this process is biased towards the training data and so may lead to model over-fitting since these are the only data that the model has seen, and no independent data has been tested. A more robust approach is to use three data sets (these are defined as following as these terms sometimes vary between laboratories):

Training set: refers to the *x*-data and *y*-data pairs used to construct a model. *Validation set*: refers to the *x*-data and *y*-data pairs used to validate model construction. This is used during training where the *y*-data predicted and *y*-data known are compared. *Test set*: refers to the *x*-data used to test the model. These *x*-data are only used after the model has been constructed with the training and validation data sets.

The use of three data sets as detailed above allows the independent assessment of the model that has been constructed using new hitherto unseen metabolite data.

When supervised analyses are used, or pre-processing optimisation employed, the above validation approach must be conducted and be included in the report.

Reporting structure:

The exact details of how the metabolite data were objectively split into training, validation and test sets must be given.

Metric used for choosing the number of latent variables, number of iterations or populations must be given based on the above three data sets.

Software

One for discussion – I have not included any at this stage as we need to be fully inclusive and the analysis strategy is more important than software X vs. software Y.

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